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- (71) Applicant: DAINIPPON INK AND CHEMICALS, INC. Itabashi-ku, Tokyo 174-8520 (JP)
- (72) Inventors:
 TANAKA, Hisakazu
 Osaka 595-0015 (JP)

- KATO, Toshiya
 Osaka 591-8031 (JP)
- IDEGUCHI, Shigeki Osaka 595-0026 (JP)
- ISHIZU, Hideyuki
 Osaka 594-0073 (JP)
- HASEGAWA, Yoshiki
 Hyogo 662-0037 (JP)
- (74) Representative: Albrecht, Thomas, Dr. Kraus & Welsert, Thomas-Wimmer-Ring 15 80539 München (DE)
- (54) WATER-ABSORBING MATERIAL AND ABSORBENT ARTICLE EMPLOYING THE SAME
- (57) The water absorbent material of the present invention is composed of a copolymer of an anhydropolymin acid having at least one ethylenically unsaturated double bond an amolecule, a water-soluble monomer having an ethylenically unsaturated double bond and polysaccharides, and has high water absorption ratio

and high water absorption rate in pure water or water having a low ion content and also has high absorption properties for high concentration salt-containing solutions.

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Description

TECHNICAL FIELD

- [0001] The present invention relates to a novel and useful water absorbent material and an absorbent article which utilities such a material. In particular, the present invention relates to a water absorbent material and an absorbent article for use in the absorption of solutions containing a high concentration of salls, such as sea water, calcium chloride deliquescent solutions, blood, and other body fluids (such as urine and sweat).
- [0002] A water absorbent material of the present invention can be widely used in fields such as sanitary products and household articles such as disposable dispers, pade, and sanitary naphics, water sealing materials, soll orditioners, anti-condensation coatings, water-storing materials for use in agriculture and horticulture, and water swelling schools.

BACKGROUND ART

[0003] Conventional water absorbent materials include hydrolysates of graft polymers of starch and acrylonitrile, and partially neutralized products of crosslinked polyacrylic acid. Furthermore, examples of water absorbent polyamino acid based resins obtained by hydrolyzing a partially crosslinked product of polyamino polyaspartate have also been disclosed in Japanese Unexamined Pätent Application, First Publication, No. Hol 9-180840. However, water absorbent materials produced from polyamino acid based resins display insufficient gel strength. Furthermore, although water absorbent materials produced from acrylic resins are typically capable of absorbing between several hundred and several thousand times their own weight of fresh water, this water absorbent materials produced from the strength of the site of th

[0004] As the water absorbent material having an enhanced water absorption capability for water containing salts, for example, the following suggestions have been made wherein an ionic water absorbent material having a small salt resistance is used in combination with a nonionic water absorbent material having a large salt resistance.

- [0005] (1) A water swelling polymer comprising a copolymer of an ethylenically unsaturated monomer with a carboxyl group and a base thereof, and a polyoxyalkylene glycol allyl ether with a hydrophobic group at one terminal (Japanese Unexamined Patent Application, First Publication, No. Sho 62-2740B); and (2) a water absorbent polymer comprising a copolymer of an ethylene based unsaturated monomer with a carboxyl group and an associated base, and an alkyl-polyoxyalkylene glycol monofinethjacrylate with an alkyl group at one terminal (Japanese Unexamined Patent Application, First Publication, No. Hai 3-93815).
- 39 [006] However, in the conventional water absorbent materials described above, although the important characteristics such as the water absorption capability (volume or water absorption, and vater absorption) and the satt tolerance are improved to some extent, the improvement is not always sufficient.
 (00007) Eurithermore, in order to solve the problems described above, water absorbent materials comortising a copol
 - ymer of either a suffoalvyf (meth)acrylate or an acrylamide (Japanese Unexamined Patent Application, First Publication, No. Hel 10-871714) or a copolymer of a nonlonic monomer and acrylic acid (Japanese Unexamined Patent Application, First Publication, No. Hel 91-43210) have also been proposed. However, although these water absorbent materials offer an improved water absorption of water which contains salts, the water absorption for pure water or water with only small amounts of lons actually decreases, and the initial water absorption speed is also show.
- [0008] In addition, as an example of a water absorbent material comprising a combination of a polyamin odd and a copolymer comprising a polyacrylic acid, the water absorbing agent composition comprising a polyamino acid and a crosslinked polyacrylate polymer as the main constituents is disclosed in Japanese Unoxamined Patent Application, First Publication, No. Hel 7-310021. However, this water absorbent material exhibits little water absorbtion capitally, and even when the surface of the water absorbtent replaced using a surface crosslinking agent, the gel strendt disclosurs on immovement.
- 0 (009) An object of the present invention is to provide a water absorbent material which displays superior water absorption of solutions with high concentrations of salts such as sea water, colcium chioride deliquescent solutions, blood, and other body fluids (such as urine and sweat) and further suffers no deterioration in the water absorption ratio or the water absorption or the water absorption ratio or the water absorption solutions.

DISCLOSURE OF INVENTION

[0010] As a result of intensive investigations aimed at resolving the problems described above, the present inventors have found that a water absorbent material comprising a copolymer of an anhydropolyamino acid having at least one

ethylenically unsaturated double bond within each molecule (A), and a water soluble monomer having an ethylenically unsaturated double bond (B) displayed a high level of water absorption capability relative to water incorporating a high concentration of salts, and as a result were able to complete the present invention.

[0011] In other words, the present invertion provides a water absorbent material comprising a copolymer of an anhydropolyamino acid having at least one ethylenically unsaturated double bond within each molecule (A), and a water soluble monemer having an ethylenically unsaturated double bond (B).

[0012] The present invention also provides a water absorbent material comprising a copolymer of an anhydropolyamino acid having at least one ethylenically unsaturated double bond within each molecule (A), a water soluble monomer having an ethylene based unsaturated double bond (B), and a polysaccharide (C).

[0013] In addition, the present invention also provides an absorbent article comprising a liquid-permeable sheet, a liquid-impremable sheet, and an absorber comprising a water absorbent material and a fiber material arranged between the liquid-permeable sheet and the liquid-impermeable sheet, wherein the water absorbent material is a water absorbent material comprising a coopymer of an anhydropolyamino acid having at least one ethylenically unsaturated double bond in a molecule (A) and a water soluble monomer having an ethylenically unsaturated double bond called to the short garden and the soluble monomer having an ethylenically unsaturated double bond called the soluble monomer having an ethylenically unsaturated double bond called the soluble monomer having an ethylenically unsaturated abusing the soluble monomer having an ethylenically unsaturated.

[0014] In addition, the present invention also provides an absorbent article comprising a liquid-permeable sheet, a liquid-impermeable sheet, and an absorber comprising a water absorbent material and a fiber material stranged between the liquid-permeable sheet, wherein the water absorbent material is a water absorbent material comprising on copolymer of an anhydropolyamino acid having at least one ethylenically unsaturated doubtle bond in a molecule (A), a water soluble monomer having an ethylenically unsaturated doubte bond on a molecule (A), a water soluble monomer having an ethylenically unsaturated doubte bond on a molecule (A), a water soluble monomer having an ethylenically unsaturated doubte bond (B), and polyacocharities (Ci).

BEST MODE FOR CARRYING OUT THE INVENTION

from any of the above methods,

Discount of

[0015] A detailed description of the water absorbent material according to the present invention is as follows,

[0016] First, the water absorbent material comprising a copolymer of an anhydropolyamino acid having at least one ethylenically unsaturated double bond in a molecule (a) (hereinafter referred to as a polymerizable anhydropolyamino acid (A)) and a water soluble monomer having an ethylenically unsaturated double bond (B) (hereinafter referred to as a water soluble polymerizable monomer (B) will be excellented.

[0017] Examples of the polymerizable anhydropolyamino acid (A) include those prepared by reacting an anhydropolyamino acid having no ethylamically unsaturated double bond in a molecule (A-1) (hereinafter referred to as an anhydropolyamino acid (A-1)) with a compound which has an ethylenically unsaturated double bond and a functional group having reactivity with the anhydropolyamino acid in a molecule (A-2) (hereinafter referred to as a polymerizable compound (A-2)) or those prepared by the polycondensation reaction of maleic amhydride, fumanic anhydride or malic anhydride and armmonia with heating.

39 [0018] Examples of the anhydropolyamino acid (A-1) include anhydrides of polyaspartic acid and polyglutamic acid. Among these anhydropolyamino acids, polysuccinimide as an anhydride of polyaspartic acid is preferred in view of industrial availability. These compounds may have a linear structure or a branched structure.

[0019] In addition, a basic skeleton of the anhydropolyamino acid (A-1) may contain a unit of an amino acid other than glutamic acid and aspartic acid.

(9020] Examples of the unit of the amino acid other than glutamic acid and aspartic acid include units of aliphatic oramino acid such as glycine, alianine, valine, leucine, isoloucine, serine, threonine, asparagine, glutamine, lysine, ornithine, cysteine, cystine, methionine, proline, hydroxyproline, or arginine, aromatic oramino acid such as tyrosine, phenylalanine, tryptochan, or histidine; cramino acid whose side chain functional group is substituted; aminocarboxylic acid such as Palanine or "**, aminobutyric acid; dispeptide (dimer) such as glycyl-glyche or aspartyl-phenylalanine; and tripeptide (trimer) such as glutarhione. These amino acids may be optically acid we substances (L-isomer, D-isomer) or racemic modifications. These amino acid units may oxist in the form of a random copolymer or a block copolymer after being ocombined with ottuamine acid or aspartic acid.

[9021] There are no particular restrictions on the method of producing the drorementioned enhydrosolyamino acid (A-1). Examples of suitable production methods include (1) heating D/L-aspartic acid and performing a dehydration condensation; (2) heating D/L-aspartic acid and performing a dehydration condensation in the presence of a catalyst such as phosphoric acid, (3) heating D/L-aspartic acid in a suitable solvent and in the presence of a catalyst such as phosphoric acid. and performing a dehydration condensation, (4) heating and reacting make an hydrogolyamino acid via emalemide or a maleamic acid in the memoria, and forming the anhydropolyamino acid via emalemide or a maleamic acid in and the presence of a catalyst such as a maleamic acid, and then producing the anhydropolyamino acid by further reaction in the presence of a catalyst such as phosphoric acid. A water absorbert material of the present invention can utilize an anhydropolyamino acid obtained

[0022] There are no particular restrictions on the polymerizable compound (A-2), although from the viewpoint of

reactivity, compounds represented by the general formula [i] shown below are preferred.

$$R^{1-Q}$$
— C — $C=CH_2$...[I]

- (wherein, R¹ represents at least one type of functional group selected from a group consisting of an amino group, an epoxy group, a carboxyl group, a carbodilmide group, an oxazoline group, an imino group and an isocyanate group, Q represents an alkylene group of 1 to 10 carbon atoms, and R² represents a hydrogen atom or an alkyl group of 1 to 4 carbon atoms.)
- [0023] Examples of compounds represented by the by the aforementioned general formula [I] include glycidyl methios acrylate, glycidyl acrylate, acrylic acid, methacrylic acid, 2-methacryloyloxyethyl isocyanate, and 2-isocyanatomethyl acrylate
- [0024] Examples of methods of reacting the anhydropolyamino acid (A-1) and the polymerizable compound (A-2) include (1) adding the polymerizable compound (A-2) directly to a powdered sample of the anhydropolyamino acid (A-1) and mixing; (2) dissolving the anhydropolyamino acid (A-1) in an aprotic organic solvent such as dimethy formamide, dimethyl acetamide, M-methyl pyrroidone, N.N-dimethyl middazolinone, dimethyl suffoxide, or suffoliane, and then adding and mixing the polymerizable compound (A-2); (3) dispersing the anhydropolyamino acid (A-1) in a solvent use as cyclohexane, hoptane, methanol, or acetone in which the anhydropolyamino acid (A-1) is insoluble, and then adding the polymerizable compound (A-2) to the dispersion and mixing, and (4) hydrolyang the anhydropolyamino acid (A-1) by adding an alkali aqueous solution to the anhydropolyamino acid (A-1), and then adding the polymerizable compound (A-2) to the dispersion and mixing.
- [0025] The amount of the polymetrable compound (A-2) used should preferably be within a range from 0.8 to 3.0 mol, and more preferably be in a range from 0.9 to 2.0 mol per 1 mol of the anhydropolyamino acid (A-1). When the amount of the polymerizable compound (A-2) relative to the anhydropolyamino acid (A-1) is within a range from 0.8 to 3.0 mol, the amount of an unreacted substance can be reduced and formation of an insoluble substance and coloration due to secondary reaction can be prevented, thereby makin it is usossible to inhibit factors which exact an adverse in-
- [0026] When the amount of the polymerizable compound (A-2) is within the above range, the amount of an unreacted substance can be reduced and formation of an insoluble substance and coloration due to the crosslinking reaction as the secondary reaction can be prevented, thereby making it possible to Inhibit factors which exert an adverse influence on the product.
- [0027] Conditions of the reaction between the anhydrop obyamino acid (A-1) and the polymerizable compound (A-2) are not specifically limited, but the reaction is preferably conducted at a temperature within a range from 20 to 150°C. The reaction time is preferably two hours or less.
- [0028] The molecular weight of the resulting polymerizable annydropolyamino acid (A) to be used in the present invention is proferably 500 or more in terms of weight, average molecular weight (horeinafter referred to as Mw), and more preferably 1000 or more. When the molecular weight is 500 or more, a water absorbent material having sufficiently enhanced water absorption properties to salts-containing water as the object of the present invention can be obtained. [0029] A portion or all of the annydropolyamino acid (A) is preferably hydroyca. An acidic amino acid nosticulo water absorption properties for salts-containing water, is formed by hydrolysis. The hydrolysis amethod is as described below.
 - [0030] As the water-soluble polymerizable monomer (S) used in the present invention, for example, there can be used lonic monomer such as (moth) apply del and/or its alkel in metal sett, lakel learth metal sati, read monitor used lonic monomer such as (moth) applyamide. N.N-dimethylacrylamide, 2-hydroxyethyl (meth)acrylate; or N-methylac (meth)acrylate; and long court on the substitution of the substitu

fluence on the product.

group. These water-soluble polymerizable monomers can be used alone or in combination.

[0031] Examples of the alkali metal salt of (meth)acrylic acid and sulfonic acid include sodium salt, potassium salt, illihium salt, and rubidum salt. Among those alkali salts, sodium salt or potassium salt is preferred in view of the performances of the resulting polymer, industrial availability, and safety.

- [0032] As used herein, the term "(meth)acrylic" means "acrylic" and 'methacrylic". In these water-soluble polymerizable monomers, (meth)acrylic acid and/or its alkali metal sall, ammonium sall and (meth)acrylamide are preferred in view of water absoruble no procerties.
- [0033] Among these water-soluble polymetrizable monomers, a water-soluble polymerizable monomer having an ethylenically uneaturated double bond such as alkali earth metal salt or ammonium salt and a suffonia acid group and/ or a sulfoniate group (hereinafter referred to as a sulfonia acid group-containing polymerizable monomer) is preferably used because the water absorption properties as well as the water absorption ratio and the initial water absorption rate of pure water and water having a low ion content are not followered.
- [0034] In addition to the water-soluble polymerizable monomer (B), other hydrophobic monomers having an ethylenically unsaturated double bond can also be used as far as water absorption performances of the copolymer constitution the water absorbent material of the present invention are not impaired.
 - [0035] Examples of the other hydrophobic monomer having an ethylenically unsaturated double bond include various acrylic acid esters such as methyl acrylate, eithyl acrylate, bubyl acrylate, and cyclobacyl acrylate; vanous methacrylic ecid esters such as methyl methacrylate, bubyl methacrylate, is-ob-utyl methacrylate, bubyl methacrylate, bubyl methacrylate, cyclobacyl methacrylate, acrylate and benzyl methacrylate, and various diesters of unsaturated carboxylic acid and alcohol, such as dimethyl fumarate, dibryl fumarate, diotryl fumarate, diotryl fumarate, diotryl fumarate, diotryl faconate, didiryl faconate, diotryl faconate, diotr
- These hydrophobic monomers can be used alone or in combination.

 [0036] The amount of the water-soluble polymerizable monomer (8) is usually within a range from 0.1/1 to 100/1 in terms of a weight ratio (weter-soluble polymerizable monomer (8) polymerizable monomer (8) is within the above range, a reably from 1/1 to 50/1. When the amount of the water-soluble polymerizable monomer (8) is within the above range, a water absorbent material having excellent water-absorption properties for salts-containing water can be obtained.

 [0037] The copolymer of the water absorbent material of the present invention preferably comprises gel particles having a crosslinked structure introduced therein to enhance the strength of the copolymer. When the quantity of the
- crosslinked structure in the copolymer increases, it becomes possible to enhance the strength of the copolymer. On to the other hand, when the quantity of the crosslinked structure decreases, it becomes possible to enhance water absorption properties. Therefore, the gel strength and water absorption properties of the copolymer can be adjusted by appropriately controlling the crosslinked structure.
 - [0038]. The get strongth of the water absorbent material of the present invention is preferably 0.1 g/cm² or more, the upper limitation being a numerical value where desired water absorption properties can be obtained. The get strength is based on the numerical value measured by the "method of measuring the get strength" described heroinafter.
 - [0039] Examples of the method of preparing gel particles include (1) a method of irradiating the copolymer used in the water absorbent material of the present invention with active radiation such as electron beams, radiation or the like and (2) a method of using a crosslinking agent.
- [0040] Examples of the crosslinking agent include crosslinkable monomers having at least two ethylenically unsaturated double bonds, crosslinkable monomer having at least two reactive groups, and crosslinking agents other than these crosslinking agents.
 - [0041] As the crosslinkable monomer having at least two ethylenically unsaturated double bonds, any monomer having two or more ethylenically unsaturated double bonds and examples thereof include N.N'-methylenebles (meth) acrylamide, (poly)ethylene glycol di(meth)acrylate, (poly)propylene glycol di(meth)acrylate, trimethylopropane tri (meth)acrylate, glycorin tarylate marylate matylate methacrylate, ethylene oxide-modified trimethylolpropane trimethylogrivate, potaneophiriol (tetra(meth)acrylate, dentacrythrtol hexa(meth) acrylate, triallyl cyanurate, triallyl sicyanurate, triallyl phosphate, triallylamine, poly(meth)allyloxyalcane, (poly)ethylene glycol, propylene glycol, propylene glycol, glycerin, pentacrythrtol, ethylenediamic, polystylenemine, and glyciol/ (meth)acrylate, despinediamic, and glycol) (meth)acrylate).
- 260 [0042] Examples of the crosslinkable monomer having at least two reactive groups include polyhydric alcohol such as attylence glycol, diethylene glycol, tiethylene glycol, polyentylene glycol, 1,4-butanediol, 1,5-pentanediol, 1,6-hexanediol, neopentyl alcohol, diethanedimine, tridelinhanelmine, polypropylene glycol, polyvinyl alcohol, pentaerythritol, sorbitol, sorbitan, glucose, manntiol, mannitane, sucrose, or glucose, polyprodyl enter, or glycorin triglycioly ether, or glycorin triglycioly ether, as entylene glycol gycidy ether, or glycorin triglycioly ether, as electronic polypropylene glycoly compound such as epichlorohydrin or cr-methylchlorohydrin, polyaledryde such as cylinalidathyd or gylosen.
- loepoxy compound such as epichlorohydrin or α-methylchlorohydrin, polyaldehydd such as glutalaiddhydd or glyoxazol, polyamines such as ethylenediamine; and hydroxide, hailde, cartonate, oxide, borate (e.g. borax, etc.) or polyvalent metal compound (e.g. aluminum isopropylate, etc.) of a metal of the group IIA, IIIB and VIII of the Periodic Table, such as calcium hydroxide, calcium chloride, calcium carbonate, calcium oxide, borax magnesium chloride, magnesium

oxide, aluminum chloride, zinc chloride, or nickel chloride.

[0043] These compounds can be used alone or in combination, taking reactivity into consideration.

[0044] The amount of the crosslinkable monomer having at least two ethylenically unsaturated double bonds or the crosslinkable monomer having at least two reactive groups is preferably within a range from 0.005 to 2 mol%, and more preferably from 0.01 to 1 mol%, based on the water-soluble polymerizable monomer (B). When the amount is within a range from 0.005 to 2 mol%, a water absorbent material having a good balance between the water absorption properties and oel strendth; can be obtained.

[0045] Examples of the crosslinking agent include diglycidyl ether compound, haloepoxy compound, polyamine compound, and iscovanate compound.

[0046] Examples of the diglycidyl ether compound include ethylene glycol diglycidyl ether, propylene glycol diglycidyl ether, and gycorin-1,3-diglycidyl ether. Examples of the halocpoxy compound include ephthorohydrin and β-mathyl-epitholrohydrin. Examples of the polyarine compound include chain alighatic polyarine such as chrylenediamine, dicitylenetriamine, trightylenetetramine, tetraethylenepentamine, pentaethylenehexamine, hexamethylenediamine, or polyather polyamine; cyclic alighatic polyamine such as menthenediamine, pentaethylenediamine, or ble(4-amiloropsy)12-4,8,10-letraevanepyrol[5,5]-undecane; aromatic polyamine such as m-vylenediamine or p-xylenediamine; polyamides obtained from dimer acid and an alighatic polyamine; and basic amino acid such as lysine.

[0047] Examples of the isocyanate compound include tolylene diisocyanate (TDI), phenylene diisocyanate (PPDI), diphenylmethane diisocyanate (MDI, hydrogenated MDI, polymeric MDI. Iolicinie diisocyanate (TDDI), hexamethylene diisocyanate (HDI), sophornee diisocyanate (XDI), bylene diisocyanate (TDDI), bylene diisocyanate (TDDI), tertamethylene yasylene diisocyanate (TMXDI), triphenylmethane triisocyanate, trisicocyanate, phonylmethylene triisocyanate, bylene ester triisocyanate, and urethane-modified compounds thereof, alophanate-modified compound, burd-modified compound, socyanurate-modified compound, contramed the compound that the contramed that the cont

[00:48] The crosslinking agent is preferably used as long as water absorption properties of the water absorbent material are not impaired, and the amount is usually within a range from 0.1 to 60 mol%, and preferably from 1 to 50 mol%, based on the inflied rind of the antivoropolyamino acid.

[0049] The crosslinkable monomer is used in the reaction between the polymerizable anhydropolyamino acid (A) and the water-soluble polymerizable monomer (B). The crosslinking agent is preferably added during or after the reaction between the polymerizable anhydropolyamino acid (A) and the water-soluble polymerizable monomer (B).

[0050] The water absorbent material comprising a copolymer of the polymerizable anhydropolyamino acid (A), the water-soluble polymerizable monomer (B) and polysaccharides (C) will now be described.

[0051] In the copolymer constituting the water absorbent material of the present invention, a molety having high nonionicity is introduced by using polysaccharides (C) as a copolymer component, thereby making it possible to further enhance water absoration properties to salts-containing water.

[0052] Examples of the method of copolymertzing polysaccharides (C) with the polymertzable anhydropolyamino acid (A) and the water-soluble polymerzable monomer (S) include (1) a method of integropening polysaccharides in the presence of an azo catalyst, thereby to activate carbon atoms to which hydroxyl groups are attached and to graft-copolymerize carbon atoms with unsaturated double bonds and (2) a method of using a crosslinking agent capable of reacting with the respective functional groups of the polymerizable anhydropolyamino acid (A), the water-soluble polymerizable anhydropolyamino acid (A), the water-soluble polymerizable anhydropolyamino acid (A), the water-soluble polymerizable anhydropolyamino acid (A) the water-soluble polymerizable anhydropolyamino acid (A).

obtained by any method can be used

[0053] Examples of polysaccharides (C) include starch, cellulose, and alginic acid.

45 (0054) Examples of the starch include starch tembers, and agrin, activities of plants, estarch-containing substances, and modified substances end/or amylopectin originating in natural substances or plants, starch-containing substances, and modified substances thereof. Specific examples thereof include potal starch, come starch, seep starch with starch seep starch and acrylic acid starch, end starch starch into destrin or sytrone, those obtained by greating starch with fatty acid, and those obtained by conversion starch into destrin or exidation, progektatinization treatment, etherification, esterification or crosslinking of starch. In addition, a structure-modified starch obtained by heating by drous starch to a temperature higher than its glass transition imperature and multing plorit (described in EPA-S2505) is also included. Furthermore, polysaccharides such as guar gum, chitin, chitosan, cellulose, algrine acid, and agar can be used.

[0055] Examples of the cellulose include celluloses obtained from wood materials, leaves, stems, basts and seed fibers; and processed celluloses such as allyl-etherified cellulose, organic acid-esterfiled cellulose, carboxymethylated cellulose, cellulose oxide, and hydroxyatip-therified cellulose.

[0056] The amount of polysaccharides (C) is usually 10/1 or less in terms of a weight ratio [polysaccharides (C) /

polymerizable anhydropolyamino acid (A)], and preferably 5/1 or less. By using polysaccharides (C) within the above range, it becomes possible to obtain the effect that the water absorbent material has non-lonicity.

[0057] Similar to the water absorbent material, the monomer having at least two ethylenically unsaturated double bonds or the crosslinkable monomer having at least two reactive groups and the curing agent can be used as the monomer component of the copolymer.

[0058] The method of preparing the copolymer constituting the water absorbent material of the present invention will now be described.

[0059] The method of preparing the copolymer can be conducted by a well-known method. That is, there can be used any method such as (1) a method of charging a polymerizable anhydropolyamino scid. (A) and a water-soluble polymerizable monomer [8] in a reaction vessel at a time, mixing them and reacting the mixture or (2) a method of initiating the reaction of one component and adding the other component dropvies. In the present invention, this is not

[0060] It is preferred in view of uniform reaction that polysaccharides (C) be previously dissolved or swollen and dispersed in a system before the reaction.

folds1 in the case of reacting the polymerizable anhydropolyamino acid (A) with the water-soluble polymerizable monomer (R) although a method of polymerizable monomer (R) although a method of polymerizable with readiation, electron beams, or ultraviolet rays, cambe employed, a polymerizable method using a radical polymerizable initiator is industrially preferred. Specific sourples of the radical polymerizable initiator include inorganic peroxide such as hydrogen peroxide, ammonium persultate, potassium persultate, regrain peroxide such as benzylip proxide, di-hobylipperoxide, currently proxide, di-hobylipperoxide, currently proxide, di-hobylipperoxide, di

specifically limited.

droxy peroxide, succinic acid peroxide, or di(2-ethoxyethy) peroxycarbonate; azo compound such as azobisisobutyronitrile, azobisyanovaleric acid, or 2.2-azobis(2-aminodipropane) hydrochloride; and redox catalyst (mado of a combination of a reducing agent such as sulfite or hyposulfite of alkali metal, armonium sulfite, armonium bleutifite or ascorbic acid and an oxidizing agent such as persulfate of alkali metal, armonium persulfate, or peroxide. These radical polymerization initiators may be used alone or in combination.

25 [0062] The amount of the radical polymerization initiator is usually within a range from 0.0001 to 5% by weight, and preferably from 0.0005 to 1% by weight, based on the total amount of the water-soluble polymeriable monomer (S) and the crosslinking agent.

[0063] In the polymerization reaction, hydrophilic polymers such as polyacrylic acid or its salt, crosslinked substance thereof, polydrinj pyrrolidone, and polywinj alcohol; chain transfer agents such as hypophosphorous acid and alkylmercaptars, surfactants; and blowing agents such as carbonate, dry ice, and aze compound can be added.

[0064] The polymerization reaction may be conducted in an aqueous solution, a solvent and a suspension, and is not specifically limited.

[0065] In the case in which the polymerization reaction is conducted in an aqueous solution, it is preferred to previously hydrolyze the polymerizable anhydropolyamino acid (A).

5 [0066] The hydrolysis reaction is conducted by an aqueous solution of an alkali metal compound and/or an alkali earth metal compound under the conditions that the reaction temperature is usually within a range from 0 to 100°C, and preferably from 20 to 50°C. The reaction time is not specifically limited, but is usually 20 hours or less, preferably 10 hours or less, and particularly preferably 2 hours or less, in view of the productivity.

[0067] Typical examples of the alkali metal compound or the alkali earth metal compound include hydroxide and or carbonate of the alkali metal and alkali earth metal. Specific examples thereof include lithium hydroxide, sodium hydroxide, because hydroxide, potalem metabonate, sodium carbonate, potassium carbonate, magnesium carbonate, and calcium carbonate. Generally, an aqueous 0.1-40% wt% solution of sodium hydroxide in the potalem hydroxide is used. The amount of the alkali compound added is an amount corresponding to 0.4 to 1 mlo arm of the inide ring group.

45 [0068] In the case in which the polymerization reaction is conducted in a solvent, the components are dissolved in the solvent. Examples of the solvent include aprotic organic solvents such as dimetriylformamide, dimetriylacetamide, N-metriylptyrolidone, N,N-dimetriylimidasciolone, dimetriyli sulfoxide, and sulfolane.

[0069] In the case in which the polymerization reaction is conducted in a suspension, a reverse phase suspension polymerization method can be used.

[0070] The reverse phase suspension polymerization method will now be described.

[0071] Examples of the reverse phase suspension polymerization method include (1) a method of reverse phase suspension polymerization of an aqueous mixed solution of a polymerizatio anhydropolymino acid (A) and a water-soluble polymerizatio monomer (B) in a hydrophobic solvent containing a water-in-oil type thereinafire referred to as W/O type) surfactant in the presence of a crosslinking agent, using a water-soluble radical polymerization initiator; (2) a method of initiating reverse phase suspension polymerization of a water-soluble polymerization monomer (B) in a hydrophobic solvent containing a W/O type surfactant in the presence of a consisting agent is not a water-soluble hydrophobic solvent containing a w/O type surfactant in the presence of a consisting agent using a water-soluble

hydrophobic solvent containing a W/O type surfactant in the presence of a crosslinking agent, using a water-soluble radical polymerization initiator, and further conducting reverse phase suspension polymerization by adding dropwise an adueous solution of a polymerizable anhydropolyamino acid (A); and (3) a method of conducting first-stage reverse

phase suspension polymerization of an aqueous solution of a water-soluble polymerizable monomer (3) in a hydrophobic solvent containing a WO type surfactant in the presence of a crosslinking agent, using a water-soluble radical polymerization initiator, and further conducting reverse phase suspension polymerization by adding a mixed solution of a polymerizable anhydropolyamino acid (A) and the water-soluble polymerizable monomer (B). In the case of the copolymer constituting the water absorbent material of the present invention, any reverse phase suspension polymerization method can be used.

[0072] This reverse phase suspension polymerization method is a preferred method because a bead-like water absorbent material capable of being easily ground can be obtained by using a surfactant.

[0073] Specific method of preparing a copolymer by the reverse phase suspension polymerization method using a sulfonic acid group-containing polymeriable monomer as the water-soluble polymerizable monomer (B) will now be

[0074] An aqueous solution of a sulfonic acid group-containing polymerizable monomor (B) is prepared by adding and dissolving acroselnifing agent and reactional polymerizablor initiation and increases and a varie-recoluble chain transfer agent such as thiols, thiol acids, secondary alcohols, amines or hypophosphites in an aqueous solution containing a previously neutralized sulfonic acid group-containing polymerizable monomer and the other monomer having dibylenically unsaturated double bonds, and then the resulting solution is deserted by introducing an inert gas such as nitrogen. In a polymerization apparatus, a surfactant is charged in a hydrophobic solvent and is optionally dissolved by heating, and then aeration is conducted by introducing an introgen gas into the apparatus. The aqueous solution of the sulfonic acid group-containing polymerizable polymer is poured into the apparatus and temperature rising is initiated under string. During temperature rising, the aqueous solution is converted into water droplets, which are suspended while being dispersed in the hydrophobic solvent. With the temperature ning, heat is generated and the polymerization is initiated.

[0075] The method of adding the polymerizable anhydrocolyamino acid (A) is not specifically limited, but includes, for example, (1) a method of previously mixing an aqueous solution of a previously hydrolyzed polymerizable and previously mixing an approach solution of a sulfance acid group-containing polymerizable monomer (B), (2) a method of simultaneously pouring an aqueous solution of a sulfance acid group-containing polymerizable monomer (B), (3) a method of pouring after the polymerization was initiated by heat generation. Among these methods, the method (4) is preferred because it can maintain the stability of the system more sulfatfactivity.

30 [0076] In the case in which the aqueous solution of the polymerizable anhydropolyamino acid (A) is added after the polymerization was initiated by heat generation, it may be added as it is.

(0077) Since there sometimes arise a problem that polymer particles agglomerate according to the amount of the polymerizable anhydropolyamine acid (A), an inter solvent containing a surfactant dissolved therein is added to an aqueous solution of the polymerizable anhydropolyamine acid (A) and the polymerizable anhydropolyamine acid (A) and the polymerizable anhydropolyamine acid (A) is dispersed in the hert solvent by stirring, and then the resulting dispersion is preferably added to a polymer solution. [0078] Although timing of pouring after heat generation is not specificably limited, pouring is conducted during the time immediately after heat generation and the time after two hours have passed since heat generation. Pouring is preferably conducted during the above timing range because the aqueous solution of the polymerizable anhydropolyamina oxid (A) can be incorporated into particles of the copolymer, which is being formed.

in the reaction system, without separating the aqueous solution.

[0079] After initiation of the polymerization, cooling is appropriately conducted according to the state of heat generation. The temperature of the polymerization reaction is preferably within a range from 60 to 100°C, and particularly preferably from 50 to 80°C.

45 [0080] The rotation number of a stirring blade of a reaction apparatus during the reaction cannot be shown unembiguously because the absolute value thereof varies depending on the kind of the stirring blade and the size of the polymerization reaction vessel, but is preferably within a range from 50 to 500 rpm in view of the polymerization safety. [0081] The suspension polymerization reaction yields a slurry mixture containing particles having an average particle diameter within a range from 10 to 300 um fiveforus cell particles/excess surfactant/hydroboble solvent).

30 [0082] The slurry mixture is converted into copolymer particles by direct dehydration according to a publicly known procedure or azeotropic dehydration with a hydrophobic solvent and optionally subjecting to a surface treatment, followed by dying and various stops such as screening.

[0083] As the W/O type surfactant used in the present invention, any surfactant can be used as long as it is soluble in a hydrophobic solvent or it has hydrophobic like a hydrophobic solvent or it has hydrophobi

[0084] The surfactant having such properties is a nonionic or anionic surfactant which generally has a HLB value within a range from 1 to 9, and preferably from 2 to 7. Specific examples of the surfactant include sorbitan fatty acid ester, polyocycerbitan fatty acid ester, poly

anhydride modified polybutadiene, maleic anhydride modified ethylene-propylene-diene-terpolymer, copolymer of α -olefin and maleic anhydride or derivative thereof, and polyoxyethylene alkyl ether phosphoric acid.

[0085] The amount of the surfactant is within a range from 0.05 to 10% by weight, and preferably from 0.1 to 1% by weight.

- [0086] As the hydrophobic solvent, any solvent can be used as long as it is basically insoluble in water and is inert with respect to the polymerization reaction. Examples thereof include allphatic hydrocarbons such as n-pentane, n-hexane, n-heptane, or n-octane; and airomatic hydrocarbons such as benzene, tolloue, or xylene. Among these solvents, n-hexane, n-heptane and cyclohexane are particularly preferred because a non-greesy water absorbent material can be obtained.
- [0087] The amount of these hydrophobic solvents is usually 0.5 to 10 times larger than that of the aqueous solution of the water-soluble polymerizable monomer (B) used in the first-stage reaction in the reverse phase suspension polymerization method (3).
 - [0088] The reaction operation may be conducted in atmospherio air, but is preferably conducted in an inert gas atmosphere to inhibit the secondary section. The reaction pressure is not specifically intrinde, but is preferably are duced pressure below normal pressure, and particularly preferably is a highly reduced pressure. Specifically, it is preferably within a range from 10 to 1.013 × 1079 B.
 - [0089] The reaction time is not specifically limited, but is usually 100 hours or less, preferably 50 hours or less, and more preferably 20 hours or less.
- [0090] The method of preparing the copolymer used in the water absorbent material of the present invention using 20 the reaction apparatus includes, for example, a preparation method using a publicly known reaction apparatus and specific examples thereof include (1) a method of polymerizing white string optionally in a twin-bowl kneader, (2) a method of cast-polymerizing in a container and (3) a method of standing-polymerizing by continuously feeding on a driving belt. The water absorbent material of the present invention can be prepared by any method, in addition to the preparation methods using the apparatus.
 - [0091] To optimize water absorption properties, about 5 to 100 mol%, preferably 65 to 80 mol% of acid groups in the water-soluble polymerizable monomer (5) is preferably neutralized with an alkali compound before or after the reaction. [0092] The alkimerizable monomer (5) is preferably a hydroxide or a carbonate of an alkali metal salt. Examples of the compound include lithium hydroxide, sodium hydroxide, backium hydroxide, magnesium hydroxide, calcium hydroxide, include lithium carbonate, and calcium carbonate, magnesium carbonate, and calcium carbonate, and calcium carbonate, and calcium carbonate.
 - [0093] The water absorbent material of the present invention varies depending on the method of the copolymerization reaction of the copolymen. In case of the reaction of an aqueous solution, the water absorbent material can be prepared by granulating hydrous get particles made of the copolymen after the reaction, and passing through a series of steps such as dryling step, grinding step, screening step, surface crosslinking treatment step and screening step.
- 5044] To dry hydrous gel particles obtained through the above steps, hydrous gel particles are preferably granulated to from granules having a predetermined particle diameter in order to enhance the drying efficiency by increasing the surface area. Hydrous gel particles are granulated simultaneously by polymerizing while stiming with a twin-bowl kneeder, or by extructing polymerized hydrous gel particles through a die using a meet grinder. Hydrous gel particles can also be granulated by a cutting mill. Although the particle diameter of granulated gel particles can be appropriately adjusted by a drier, but an average particle diameter is proferably within a range from 0.1 to 10 mm. When the average particle diameter is water shown in a diameter is less that no 1 mm, physical properties of the water absorbern material are iskely to be lowered. On the other hand, when the average particle diameter exceeds 10 mm, granulated gel particles are not easily dried and, therefore, it is not preferred.
- [0095] On granulation of hydrous gel particles made of the copolymer, gel coarse particles having an average particle diameter of larger than 10 mm and gel fine particles having an average particle diameter of smaller than 0.1 mm are likely to be formed. The gel coarse particles and gel fine particles can be reused by adding to the aqueous solution of the water-soluble polymerizable monomer (B) and polymerized gel particles after screening and recovering.
 - [0096] Gel particles granulated in the granulating step are dried in the following drying step. In the drying method, for example, a hot-air drier, air-current drier, a fluidized bed drier, a drum drier, microwave drier, a far infrared drier and a vacuum drier can be appropriately used.
 - [0037] Hydrous gel particles obtained in the drying step are ground and screened according to uses of the water assorbent material in the following grinding step and screening step to form granules having a predetermined particle size. When used in a disper, napkin or the like, the screened granules have a particle size of 1 mm or less, and preferably 0.85 mm or less. To sufficiently exert water absorption performances in the disper or sanitary napkin, fine powders having a particle diameter of 105 µm or less, preferably 122 µm or less, and more preferably 300 µm or less are preferably removed by screening. Fine powders recovered in the grinding step and screening step can be reused by adding in the polymerizing step and drying step and crying step.
 - [0098] The vicinity of the surface of the water absorbent material of the present invention can be crosslinked by using

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a surface crosslinking agent having two or more functional groups capable of reacting with functional groups of the copolymer constituting the water absorbent material.

[0099] Specifically, the water absorbent material of the present invention is mixed with the surface crosslinking agent, and then the vicinity of the surface of the water absorbent material is crosslinked by a heat treatment.

- [01001] Examples of the surface crosslinking agent include polyhydric alcohol such as diethylene glycol, propylene glycol, triethylene glycol, tetraethylene glycol, polyethylene glycol, 1,3-propanediol, dipropylene glycol, 2,3,4-trimethyl-1,3-pentanediol, polypropylene glycol, glycerin, polyglycerin, 2-butene-1,4-diol, 1,4-butanediol, 1,5-pentanediol, 1.6-hexanediol, 1,2-cyclohexanedimethanol, 1,2-cyclohexanediol, trimethlolpropane, diethanolamine, triethanolamine, polyoxypropylene, oxyethylene-oxypropylene block copolymer, pentaerythritol, or sorbitol; epoxy compound such as ethylene glycol diglycidyl ether, polyethylene glycol diglycidyl ether, glycerol polyglycidyl ether, diglycerol polyglycidyl other, polyglycerol polyglycidyl ether, propylene glycol diglycidyl ether, polypropylene glycol diglycidyl ether, or glycidol; polyhydric amine compound such as ethylenediamine, diethylenetriamine, triethylenetetramine, tetraethylenepentamine, pentaethylenehexamine, polyethylenelmine, or polyamidepolyamine; haloepoxy compound such as epichlorohydrin, epibromohydrin, or α-methylepichlorohydrin; condensate of the above polyvalent amine compound and the above haloepoxy compound; polyhydric isocyanate compound such as 2,4-tolylene diisocyanate or hexamethylene diisocyanate; polyhydric oxazoline compound such as 1,2-ethylenebisoxazoline; silane coupling agent such as y-glycidoxypropyltrimethoxysilane or y-aminopropyltrimethoxysilane; alkylene carbonate compound such as 1,3-dioxolan-2-one, 4-methyl-1,3-dioxolan-2-one, 4,5-dimethyl-1,3-dioxolan-2-one, 4,4-dimethyl-1,3-dioxolan-2-one, 4-ethyl-1,3-dioxolan-2-one, 4-hydroxymethyl-1,3-dioxolan-2-one, 1,3-dioxin-2-one, 4-methyl-1,3-dioxan-2-one, 4,6-dimethyl-1,3-dioxan-2-one, 1,3-dioxan-2-one, or 1,3-dioxoban-2-one; and polyhydric metal compound such as hydroxide or chloride of zinc, calcium, magnesium or aluminum. Among these surface crosslinking agents, polyhydric alcohol compound, epoxy compound, polyhydric amine compound, condensate of polyvalent amine compound and
- halcepoxy compound, and alkylene carbonate compound are preferred in view of the reactivity and safety. These surface crosslinking agents may be used alone or in combination.

 [10101] The amount of the surface crosslinking agent relative to the water absorbent material varies depending on the combination of the water absorbent material and the surface crosslinking agent, but is usually within a range from 0.01 to 10 parts by weight, and preferably from 0.05 to 5 parts by weight, based on 100 parts by weight of the copolymer in the dry state. By using the surface crosslinking agent in the dry state. By using the surface according agent in the amount within the above range, water absorption prop-
- erties for body fluids (aqueous liquids) such as urine, sweat or menstrual blood can be further improved.

 [10102] In the case in which the water the water absorbent material is mixed with the surface or constituting agent, water is preferably used. The amount of water varies depending on the kind, the particle size and the water obsorbent material, but is usually within a range from 0.5 to 10 parts by weight, and preferably from 0.5 a parts by weight, and to contain of the water absorbent material, thus making it ossible to form a crossilisting layer having a sufficient flickness in the vicinity of the surface.
- 35 [0103] In the case in which the water the water absorbent material is mixed with the surface crosslinking agent, a hydrophilic organic solvent include lower alcohol such as methyl alcohol, othyl alcohol, ethyl alcohol, ethyl
- 40 [0104] The amount of the hydrophilic organic solvent varies depending on the kind and particle size of the water absorbent material but is usually within a range from 0.001 to 10 parts by weight, and preferably from 0.1 to 5 parts by weight, based on 100 parts by weight of the water absorbent material.
- [0105] It is necessary that a preferred mixer used to mix the surface crosslinking agent with the water absorbent material is capable of producing a large mixing force to ensure uniform mixing. Preferred examples of the mixer include 45 a cylindrical mixer, double wall conical mixer, flight-speed stifring mixer, Vehaped mixer, thobon-shaped mixer, screw mixer, flow furnace rotary disk type mixer, air-current mixer, twin-bowl kneader, internal mixer, grinding mixer, rotary mixer, and screw extruder.
 - [0106] The temperature of the heat treatment is preferably within a range from 80 to 300°C. When the temperature is within the above range, more uniform crosslinking can be achieved, thereby making it possible to obtain a water absorbent material which is capable of reducing the amount of a soluble component eluted and has excellent water absorption properties.
 - [0107] As the apparatus for heat treatment, for example, a publicly known drier or heating furnace can be used. Specific examples thereof include groove mixing drier, rotary drier, disk drier, fluidized bed drier, air-current drier, infrared drier, and vacuum drier.
- 55 [0108] The water absorbent material of the present invention is superior in water absorption properties to pure water and salts-containing water, especially water absorption properties to salts-containing water.
 [0109] Water absorption performances can be determined by the method of esting water absorption of a high water absorbent material using a tea bag method defined in Japanese Industrial Standard (JIS K7223). In the case of eval-

uating by the leabag method, the water absorbent material of the present invention has water absorption performances of 20 times or more to ion exchange water and has water absorption performances of 5 times or more to a physiological saline (aqueue) 9.9% sodium chloride solution).

- [0110] The water absorbent material of the present invention can be applied to all purposes that have conventionally 5 been known. The water absorbent material can be used in the fields, for example, sanitary field such as sanitary products (e.g. diapers, sanitary napkins, etc.), medical field such as poulleas, civil engineering and construction field such as sludge gelling agents, food field, industrial field, and agricultural/horticultural field such as soil conditioners and water-storing materials and its utility value is remarkably great. Other purposes include water-swelling rubber prepared by incorporating a rubber into the water absorbent material.
 - [0111] The absorbent article of the present invention will now be described.
 - [0112] The liquid-permeable sheet constituting the absorbent article of the present invention means a sheet made of a material which is permeable to an aqueoue liquid, and oxamples thereof include nonwoven fabric, woven fabric, and synthetic films made of a material such as polyethylene, polypropylene, or polyamide.
- [0113] The liquid-impermeable sheet constituting the absorbent article of the present invention means a sheet made to a material which is impermeable to an aqueous liquid, and examples thereof include synthetic films made of a material such as polyethylene, polypropylene, ethylene vinyl acetate, or polyyinyl chloride, and a film made of a composite of the synthetic reals and a nonwoven fabric or a woven fabric.
 - [0114] The liquid-impermeable sheet constituting the absorbent article of the present invention can use the above water absorbent material.
- 20 [0115] Examples of the fiber material constituting the water absorbent material include hydrophobic fiber material and hydrophilic fiber materials. Among these fiber materials, hydrophilic fiber material is preferred in view of superior affinity with the solution to be absorbed. Examples of the hydrophilic fiber material include collutions fiber such as mechanical pulp or semi-chemical pulp obtained from wood materials; artificial cellulose fiber such as rayon or acetate; and fiber material obtained by hydrophilization of themoplastic fiber. The fiber material may have a fibrous shape, or may be formed into a sheet such as issue paper or pulp mat.
- [0116] The absorbent article of the present invention comprises a liquid-permeable sheet, all quid-impermeable sheet, and an absorber comprising the water absorbert material, and a fiber material arranged between the liquid-permeable sheet and the liquid-impermeable sheet, and has a structure such that the absorber is supported inside.
 Specific examples of the method of producing the absorbert article include a method of sandwiching the absorber between the liquid-permeable sheet and the liquid-impermeable sheet and bonding an outer peripheral portion of the liquid-permeable sheet and the liquid-impermeable sheet and sheet and the liquid-impermeable sheet using an adhesive such as hot-melt adhesive or a bonding means such as a heat seal.
- [0117] The method of producing an absorber comprising a water absorbent material and a fiber material is not specifically limited, but includes (1) a method of forming a fiber material link or sheet and covering a water absorbent material with the sheet, (2) a method of dispersing a water absorbent material over a multi-layer fiber sheet and forming the multi-layer sheet, and (3) a method of mixing a fiber material with a water absorbent material and forming the mixture into a sheet.
- [0118] The absorbent article of the present invention is used in disposable diapers for infants, adults, and persons suffering from incontinence, and sanitary napkins. The absorbent article is particularly suited for use in disposable diapers for adults, wherein a swelling gel of the water absorbent material is drastically deteriorated due to a large amount of excretion and long time in contact with urine, among these uses because of its excellent absorbency of urine and body fullids as well as excellent urine leakage inhibition or feed.

EXAMPLES

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[0119] The following Examples further illustrate the present invention in detail, but the present invention is not limited by the Examples. In the following Examples, percentages are by weight unless otherwise specified. Various characteristics of the resin of the present invention were determined by the following methods.

- 50 [Method of measuring water absorption ratio]
 - [0120] The water absorption capability of resins obtained in the Examples of the present invention was determined in accordance with the method of testing water absorption of a high water absorbent material described in Japanese Industrial Standard JIS K7223. 0.20 g (1.00 g based on an aqueous 9.5% sodium chloride solution) of a dry resin was put in a tea bag (200 mm × 100 mm) made of a 25 mesh nytion gauze and the resin was swollen by dipping in 1000 ml of in exchange water or an aqueous 0.5% sodium chloride solution for a fixed time. After pulling out the tea bag, the solution for a fixed time. After pulling out the tea bag, the solution for a fixed time. After pulling out the tea bag was excluded. The resulting water she balank. The water

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absorption ratio W (g/g) was calculated in accordance with the following equation:

$$W = \frac{b_{60min} - c - a}{a}$$

where W is water absorption ratio (g/g), a is a weight (g) of a sample, b is a weight (g) of the sample measured after a tea bag containing the sample was dipped for a predetermined time and the solution was drained off, and c is an average value of a weight (g) of the sample measured after a tea bag containing no sample was dipped for a predetermined time and the solution was drained off.

[Method of measuring water absorption rate]

[0121] A stirrer tip was rotated at about 600 rpm in 50 g of an aqueous 0.9% sodium chloride solution in a 100 ml glass beaker and then 2 g of a sample was put in the beaker along with the Inner wall thereof. Then, a resin was added and the time (in seconds), which is required for the resin to be swollen by water absorption, thereby stopping rotation of the stirrer tip, was taken as the water absorption rate.

[Method of measuring gel strength]

[0122] 1.0 g of a crosslinked resin was allowed to absorb 100 g of pure water (water absorption by 100 times) and a dead-weight was placed on the resin after water absorption. The total weight of the dead-weight per unit area (g/ cm²) when the dead-weight placed first on the resin was taken as a gel strength.

25 [Synthetic Example 1]

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[0123] In a 2 L round bottom flask, 100 g of L-aspartic acid and 50 g of 85% phosphoric acid were charged and the reaction was conducted under reduced pressure in an oll bight at a bath temperature of 20°C for four hours using an evaporator. The resulting product (25 g) was washed several times with water and methanol to obtain polysucclin-mind has a vesult of the measurement of gel permeation chromatography (hereinafter referred to as GPC), this polysucclin-minde has a New of 20.000.

[Synthetic Example 2]

9 [0124] In a 1 L four-necked flaak coulpped with a stirrer, a thermometer, a reflux condenser, and a nitrogen gas introducing device, 8 g of mallec anhydride and 50 g of lon exchange water were added and, ster dissolving maleois anhydride by heating to 55°C, the solution was cooled to obtain a sturry of maleic anhydride. When the temperature of the system reached 55°C as a result of heating, 80 g of 25% ammonia water was added. The system was heated to 80°C and, after the reaction inset produced for three house, the resulting aqueous solution was dired to obtain a reaction intermediate. In a 2 L round bottom flask, 100 g of the reaction intermediate and 10 g of 85% phosphoric acid were charged and the reaction was conducted under reduced pressure in an oil bath at a bath temperature of 20°C for four hours using an evaporator. The resulting product was washed several times with water and methanol to obtain polysuccininde. The Mor of this polysuccinindia was measured by GPC. As a result, it was 3,000.

45 [Example 1]

[0125] In a 500 ml four-necked flask equipped with a stirrer, a thermometer, a reflux condenser and a nitrogen gas introducing device, 10 g of polysuccinimide obtained in Synnhetic Example 1 and 20 g of N,N-dimethyltomamide (hereinafter referred to as DMF) were charged and dissolved at about 60°C, and furthermore, 0.5 g of 2-methacyloyloxyethyl isocyanate was added. As a result, heat was generated. After the reaction was continued for 30 minutes, 150 g of an aqueous solution containing 3.3 g of sodum hydroxide dissolved therein was added and polysucchimide was hydrolyzed. After the temperature in the system was reduced to about 35°C, 1 g of "ALSTAR B" (progelatinized starch, manufactured by NiHON SHO(MLHIN KAKO CO, LTD), 25 g of acrytic acid and 0.25 g of diepertaerythriol hexamethacrytate were charged. After the atmosphere in the system was replaced by a nitrogen gas, 7.5 mg of 2.2-ezoblasmi-nodipropane dihydrochioride, 5 mg of acorbic acid and 57 mg of an acqueous hydrogen peroxide (35%) solution were respectively dissolved in 1 g of 10 ne exchange water. The resulting solutions were added to the above aqueous solution in this sequence and the reaction was initiated, and then the mixture was maintained at about 60°C for three hours (hereinafter referred to as 1) filst step). 2.4 g of hexamethylenedelamine was dissolved in 2 g of the oxchange water.

and the resulting solution was added to the above aqueous solution, and then the reaction was conducted. Furthermore, 10.4 g of sodium hydroxide was dissolved in 30 g of lon exchange water and the resulting solution was added, thereby to neutralize cathooyl groups originating in arvijet caed (hereinatter referred to as (1) third step). The resulting gel-like product was dried at 110°C by a vacuum drier and the resulting dry solid was ground to obtain a water absorbent material of the present invention. The evaluation results of characteristics of the resulting water absorbent material of the present invention.

[Example 2]

[0126] In the same operation and manner as in Example 1, except that the amount of acrylic acid was changed to 30 g, the amount of dipentaerythritol hoxamethacrylate was changed to 0.1 g, and the amount of sodium hydroxido for neutralization of carboxyl groups originating in acrylic acid was changed to 12.5 g, a water absorbort material of the present invention was obtained. The evaluation results of characteristics of the resulting water absorbort material are shown in Table 2.

[Example 3]

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[0127] In a 500 ml four-necked flask equipped with a stirrer, a thermometer, a reflux condenser, a dropping funnel and a nitrogen gas introducing device, 10 of prolysucchimide obtained in Synthetic Example 2 and 10 g of DMF were charged and dissolved at about 60°C, and then 0.5 g of 2-methacyloploxyeptly isocyanate was added. As a result, heat was generated. After the reaction was continued for 30 minutes, 125 g of an aqueous solution containing 2.1 g of sodium hydroxide dissolved therein was added and polysuccinimide was hydrolyzed. After the temperature in the system was reduced to about 35°C, 1 g of XLSTAR B* was charged.

[0128] In a 100 ml Erlemmeyer (flask, 25 g of acrylic acid, 0.125 g of dipentiaerythritol hexamethacrylate and 0.64 g of "THIOKALCOL.20" (laurylmoresptane, marufactured by Kao Corp.) were mixed to prepare an acrylic acid solution. 5 g of this mixed solution was added to the four-necked flask and the rest was transferred to a dropping lunnel. After the atmosphere in the system was replaced by a nitrogen gas, 7.5 mg of 2.2-azobisaminodipropane dihydrochloride, 5 mg of ascorbic acid and 57 mg of an aqueous hydrogen peroxide (35%) solution were respectively dissolved in 1 g of ion exchange water. The resulting solutions were added to the above aqueous solution (11) first step). After 10 minutes, dropwise addition of the acrylic acid solution in the dropping furnel was initiated and was completed over one hour. After one hour has passed since the completion of the reaction, 7.5 g of 2,2-azobisaminodipropane dihydrochloride, 5 mg of ascorbic acid and 57 mg of an aqueous hydrogen peroxide (35%) solution were respectively dissolved again In 1 g of ion exchange water. The resulting solutions were added to the above aqueous solution in this sequence, and then the mixture was heated to about 60°C and maintained at the same temperature for three hours (herein after referred to as (1) second selon.

[0129] 2.4 g of hexamethylenediamine was dissolved in 20 g of ion exchange water and the resulting solution was added to the above aqueous solution, and then the reaction was conducted. Furthermore, 10.4 g of sodium hydroxide was dissolved in 30 g of ion exchange water and the resulting solution was added, thereby to neutralize carboxyl groups originating in acrylic acid (1) third step). The resulting gel-like product was dried at 110°C by a vacuum drier and the resulting dry solid was ground to obtain a water absorbent material of the present invention. The evaluation results of characteristics of the resulting water absorbent material are shown in Table 2.

[Example 4]

45 [0130] In a 500 ml four-necked flask equipped with a stirrer, a thermometer, a reflux condenser, a dropping funnel and a nitrogen gas introducing device, 5 g of polysuccinimide obtained in Synthetic Example 2 and 10 g of DMF were charged and dissolved at about 60°C, and then 0.25 g of 2-methacry/olyocyachy) isocyanate was added. As a result, heat was generated. After the reaction was continued for 30 minutes, the temperature in the system was reduced to about 55°C, 1 g of a registalized starch was charged. After the atmosphere in the four-necked flask was replaced of by a nitrogen gas, 7.5 mg of 2.2-azoblasminodipropane olitydrochloride, 5 mg of asocrible add and 57 mg of an aqueous hydrogen peroxide (35°S) solution were respectively dissolved in 1 g of ion exchange water. The resulting solutions were added to the above aqueous solution, and then the mixture was heated to about 60°C and maintained for three hours ((1) first step). To the polymer obtained by this step, 25 g of expire add and 25 mg of N.N-methylenebisacylamide were added. After the atmosphere in the four-necked flask was replaced by a Introgen gas, 7.5 g of 2.2-azoblasmin-object and only of the polymer obtained by the step. 25 g of expire add and 25 mg of N.N-methylenebisacylamide were added. After the atmosphere in the four-necked flask was replaced by a Introgen gas, 7.5 g of 2.2-azoblasmin-object and the programment of the second of the above aqueous solution in this sequence, and then the mixture was heated again to about 50°C and maintained at the same temperature for three hours ((1) second to the above aqueous solution in this sequence, and then the mixture was heated again to about 50°C and maintained at the same temperature for three hours ((1) second to the above accounted to the above

resulting solution was added to the above aqueous solution, and then the reaction was conducted. Furthermore, 10.4 g of sodium hydroxide was dissolved in 30 g of ion exchange water and the resulting solution was added, thereby to neutralize carboxyl groups originating in earylic add ((1) third stop). The resulting gel-like product was dried at 11°C by a vacuum drier and the resulting dry solid was ground to obtain a water absorbent material of the present invention. The evaluation results of characteristics of the resulting water absorbent material or solven in Table 3.

[Comparative Example 1]

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[031] In a 500 ml four-necked flesk equipped with a stirrer, a thermometer, a reflux condenser and a nitrogen gas introducing device, 1 go is progelatinized starch, 25 g of anytic acid, 0.25 g of dipentacythrich lexametheropyties and 150 g of ion exchange water were charged. After the atmosphere in the system was replaced by a nitrogen gas, 7.5 mg of 12.2-aczoblasminodipropane dihydrochloride, 5 mg of ascorbic acid and 57 mg of an aqueous hydrogen peroxide (35%) solution were respectively dissolved in 1 g of ion exchange water. The resulting solutions were added in this sequence and the reaction was initiated, and then the mixture was maintained at 60°C for three hours ((1) first step). After the completion of the reaction, 10.4 g of sodium hydroxide was dissolved in 30 g of ion exchange water and the resulting solution was added, thereby to neutralize carboxyl groups originating in acrylic acid ((1) third step). The resulting gel-like product was dried at 110°C by a vacuum drier and the resulting dry solid was ground. The evaluation results of characteristics of the resulting groups originating in Table 2.

[0132] Components to be charged of Examples 1 to 4 and Comparative Example 1 are shown in Table 1.

Table

NSDOCID- - EP

Comp.		-				150.0	-	25	0.25		0.0075	0.005	0.057	3		-	.
Example 4		5	10	0.25	-		1				0.0075	0.005	0.057	3	25	0.025	0.0075
Example 3		10	10	0.5	2.1	122.9	1	25	0.125	0.04	0.0075	0.005	0.057	3			0.0075
Example 2	10		20	0.5	3.3	146.7	1	30	0.1		0.0075	0.005	0.057	3			-
Example 1	10		20	0.5	3.3	146.7	1	25	0.25		0.0075	0.005	0.057	3		-	
Components to be charged (g)	Polysuccinimide (1)	Polysuccinimide (2)	DMF	MCOEI	NaOH	lon exchange water	Pregelatinized starch	Acrylic acid	DPMA	Laurylmercaptane	ABAPHC	Ascorbic acid	35% hydrogen peroxide water	Ion exchange water	Acrylic acid	MBAA	ABAPHC
Ö								(1) First sten	,						(1) Second step		

	T	Γ	T		Γ	
ľ		ľ		1		10.4
0.005	0.057	3		71		10.4
0.005	0.057	3	,	5 .7		10.4
				4:7	7 61	17.3
				÷.,	7.01	10.4
Ascorbic acid	35% hydrogen peroxide water	Ion exchange water	Hexamethylenediamine aqueous solution	(ion exchange water: 20 g)	NaOH aqueous solution	(ion exchange water: 30 g)
			(1) Third ctan	dose numir (1)	1	

Polysuccinimide (1): polysuccinimide (Mw: 20,000) obtained in Synthetic Example 1 Polysuccinimide (2): polysuccinimide (Mw: 3,000) obtained in Synthetic Example 2 MCOEI: 2-methacryloyloxyethyl Isocyanate DPMA: dipentaerythrijo i hexamethacrylate

ABAPHC: 2,2-azobisaminodipropane dihydrochloride MBAA: N,N'-methylenebisacrylamide

Table 2

	lable	2					
	W	ater absorption ratio (g/g)					
	Ion exchange water	0.9% aqueous solution of sodium chlori-					
Example 1	230	62					
Example 2	270	70					
Example 3	237	114					
Example 4	301	107					
Comparative Example 1	400	39					

20 [Example 5]

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[0133] In a 500 ml four-necked flask equipped with a stirrer, a thermometer, a reflux condenser, a dropping funnel and a nitrogen gas introducing device, 121 g of cyclohexane was added, and then 0.9 g of sorbitan monostearate was added thereto and dissolved by heating to 50° Owhite string. The contents in the facis were cooled to 30°C.

- 5 [0144] In a 500 ml Erlemmeyer flast, 30 g of acrylic acid was charged and neutralized (75 mo%) by adding dropwise 91.8 g of an aqueous acdium hydroxide solution containing 12.5 g of sodium hydroxide dissolved therein while cooling from outside. To this solution, 21 mg of N.N-methylenebissory/amide was added and then 0.104 g of potassitum proportions of the proposition on the proposition of the propositio
- So [0135] In a separate 500 ml four-necked flask equipped with a stirrer, a thermometer, a reflux condenser, and a nitrogen gas introducing device, 7 g of polysuscinnide obtained in Synthetic Example 2 and 10 g of DMF were charged and dissolved at about 60°C, and then 0.35 g of 2-methacrylyrloxyethyl isocyranate was added. As a result, heat was generated. After the reaction was continued for 50 minutes, 10 g of an aqueous sodium hydroxide solution containing separated. After the reaction was continued above, 20.3 g of an aqueous spatially neutralized sodium acrystate solution (neutralization degree, 75 min/s) was prepared from 5 g of an aqueous spatially neutralized sodium acrystate solution (neutralization degree, 75 min/s) was prepared from 5 g of an aqueous acrylic and 5 soution and then added to the above solution. Furthermore, 1,73 g of polsastium persultate, 21 mg of N,N-methylenobisacrylamide and 7.1 mg of sodium hypochosphite menonybrate were added and dissolved. The aqueous monomer solution thus obtained was transferred to the dropping funnel and then slowly added dropwise to the polymer stury solution maintained at 20°C over 30 minutes. After the atmosphere in the flask was sufficiently replaced by nitrogen, the contents were headed to 70°C and maintained at the same temperature for three hours, and furthermore, the polymerization reaction was conducted (preinfalter referred to as (2) third state). Under reduced pressure, cytochoxane and water were emoved. The

50 [Comparative Example 2]

[0136] In a 500 ml four-necked flask equipped with a stirrer, a thermometer, a reflux condenser, a dropping funnel, and a nitrogen gas thirtoducing device, 121 g of cyclohoxane was added, and then 0.9 g of sorbitan monostearate was added thereto and dissolved by heating to 50° while stirring. The contents in the flask were cooled to 30°C. In a 500 ml Effenmeyer flask, 30 g of acrylic acid was charged and neutralized (75 mo%) by adding dropwise 91.8 g of an aqueous sodium hydroxide solution containing 12.5 g of sodium hydroxide dissolved therein while cooling from outside. To this solution, 21 mg of N,N-methylenebsachylamide was added and then 0.104 g of potassium persurlates.

evaluation results of characteristics of the resulting water absorbent material as gel-like particles are shown in Table 4.

0.0426 g of sodium hypophosphite monohydrate were added and dissolved. The resulting partially neutralized aqueous acrylic acid salt solution containing a polymerization initiator and a crosslinking agent (neutralization degree: 75 mol%) was added in the contents of the above cylindrical round bottom flask and dispersed in the cyclohexane solution containing the surfactant, and then the atmosphere in the system was sufficiently replaced by nitrogen. A bath was heated and the temperature was set to 70°C and the mixture was maintained at the same temperature for three hours, and

then the polymerization reaction was conducted (hereinafter referred to as (2) first step). [0137] In the same manner as described above, 20.3 g of an aqueous partially neutralized sodium acrylate solution (neutralization degree: 75 mol%) was prepared from 5 g of the aqueous acrylic acid solution in a 100 ml Erlenmeyer flask and was then added. Furthermore, 1.73 mg of potassium persulfate, 21 mg of N,N'-methylenebisacrylamide and 7.1 mg of sodium hypophosphite monohydrate were added and dissolved. The aqueous monomer solution thus obtained was transferred to the dropping funnel and then slowly added dropwise to the polymer slurry solution maintained at 20°C over 30 minutes. After the atmosphere in the flask was sufficiently replaced by nitrogen, the contents were heated to 70°C and maintained at the same temperature for three hours, and furthermore, the polymerization reaction was conducted ((2) third step). Under reduced pressure, cyclohexane and water were removed. The evaluation results

of characteristics of the resulting water absorbent material as gel-like particles are shown in Table 4. [0138] Components to be charged of Example 5 and Comparative Example 2 are shown in Table 3.

	Table 3	Example 5	Comp. Example 2
	Components to be charged (g)		
(2) First step	Cyclohexane	121	121
	Sorbitan monostearate	0.9	0.9
	Acrylic acid	30	30
	NaOH	12.5	12.5
	Ion exchange water	79.3	79.3
	MBAA	0.021	0.021
	Potassium persulfate	0.104	0.104
	Sodium hypophosphite monohydrate	0.0426	0.0426
(2) Second step	Polysuccinimide (2)	7	-
	DMF	10	
	MCOEI	0.35	-
	NaOH	1.44	-
	Ion exchange water	8.56	-
(2) Third step	75 mol% partially neutralized sodium acrylate	20.3	20.3
	Potassium persulfate	0.00173	0.00173
	MBAA	0.021	0.021
	Sodium hypophosphite monohydrate	0.0071	0.0071

Polysuccinimide (2): polysuccinimide (Mw: 3,000) obtained in Synthetic Example 2 MCOEI: 2-methacryloyloxyethyl isocyanate

Table 4

	Wa	ater absorption ratio (g/g)				
	Ion exchange water	0.9% aqueous solution of sodium chloric				
Example 5	615	135				
Comp. Example 2	676	67				

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[Example 6]

[D139] In a 500 ml four-necked flask equipped with a stirrer, a thermometer, a reflux condenser and a nitrogen gas introducing device, 75 g of an aqueous solution containing 20.6 g of sodium hydroxide dissolved therein was added and then 50 g of a powder of polysucchimide obtained in Synthetic Example 2 was added to prepare an aqueous solution of polysuccinimide. After the temperature was raised to 90°C, 5.0 g of glycidyl methacrytale was added and the reaction was conducted for one hour to obtain an aqueous solution containing a hydrolysate of polysuccinimide having methacrytoly (arous hittoduced therein.

[0140] In a 100 ml Erlenmeyer flask, 0,75 g of "DK ESTER F-160" (sucrose fatty acid ester (HLB=16), manufactured by DAI-(CHI KOGYO SEIYAKU CO., LTD.) was weighed and dissolved by adding 29 of cyclohexane and heating to 50°C. The resulting solution was added to 7.7 g of the aqueous solution obtained by the above operation, followed by stifring to obtain a dispersion of polysucchimide having methacry(cyt) groups introduced therein in an aqueous hydro-jyzed solution (hereinafter referred to as (3) first slep).

[0141] In a separate 500 ml four-necked flask equipped with a elitror, a thermometer, a roflux condenser, and a sitrogen again Irreducing device, 14.5 g of cyclohexane was added, and then 0.75 g of "DK ESTER F-90" was added and dissolved by heating to 50°C with stirring. The contents in the flask were cooled to 30°C. In a 500 ml Effentineyer flask, 18.4 g of sodium sulfcetylmetheroytate was added. To this solution, 18.4 g of acrylamide and 39 mg of N.Y. methyleneblsacrylamide were added and interherence, 0.05 g of armonium persuitate was added and dissolved of 30°C and the season and solution containing a polymerization initiator and a crosslinking agent, which is essentially composed of sodium sulforthylmetheropite and acrylamide as a main component, was added to the contents in a cylindrical round bottom flask and was dispersed in a siloxane solution containing a surfactant at a stirring rate of 300 pm and, at the same time, the atmosphere in the system was sufficiently replaced by infregon. Then, the polymerization reaction was linitiated by heating. After a while, heat was generated. After five minutes had passed since the peak of heat generation, her resulting dispersion of a hydrolysate of polysucchimide having methacryloid groups introduced therein in an aqueous solution was added at atime. The mixture was maintained at 80 to 85°C for three hours. After the completion of the reaction, the four-necked flask was equipped with a water dispenser and a comproind dehardering with a water dispenser and a comproind ehardering the completion of the reaction, the four-necked flask was equipped with a water dispenser and a comproind ehardering the solution was added at time.

completion of the reaction, the four-necked flask was equipped with a water dispenser and azeotropic dehydration was conducted by raising the temperature in the system to 70 to 80°C. After azeotropic dehydration was conducted until the amount of water in the system became 55% based on the solid content to be charged, the temperature in the system was cooled to about 40°C and a cyclohexane phase was separated by decantation. Subsequently, water was removed from a wet polymer by vacuum drying to obtain a water absorbent material as gel-like particles (the reinaitler referred to as (3) second step). The evaluation results of this water absorbent material are shown in Table 6.

[Comparative Example 3]

39 [0142] In a 500 ml four-necked flask equipped with a stirrer, a themmoreler, a reflux condenser, and a nitrogen gas introducing device, 146 g of cydohexare was added, and then 0.75 g of "DK ESTER F-90" was added and dissolved by healing to 50°C with stirring. The contents in the flask were cooled to 30°C. In a 500 ml Erlenmeyer flask, 184 g of sodium sulfoethylmentacytake was added. The solution is solution, 18.4 g of acrytamide and 3.9 mg of N.N'-methyleneble sterylamide were added, and furthermore, 0.05 g of ammonium presultate was added and dissolved. The resulting agent, which is assentially composed of sound to solution to containing a polymerization initiation and a crosslinking agent, which is assentially composed of sound to solution of the solution of the solution of the solution of the solution flask and was dispersed in a siloxane solution containing a surfactant at a stirring rate of 300 gm, and at the same item, be atmosphere in the system was sufficiently replaced by hirtogen. Then, the polymerization reaction is the same time, the atmosphere in the system was sufficiently replaced by hirtogen. Then, the polymerization reaction of the reaction, the four-necked flask was equipped with a water dispenser and aerostropic dehytexion was conducted by risistion the

5 the four-necked flask was equipped with a water dispenser and areotropic dehydration was conducted by reising the removerful; in the system to 70 to 80°C. After areotropic dehydration was conducted until the amount of water in the system became 35% based on the solid content to be charged, the temperature in the system was cooled to about 40°C and a cyclohoxane phase was separated by decantation. Subsequently, water was removed from a wet polymer by vacuum driying to obtain a water absorbert material as get-like particles (hereinafter referred to as (3) third step).

The evaluation results of this water absorbent material are shown in Table 6.

[Example 7]

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[0143] In a 800 ml Efnermeyer flask, 16.5 g of 2-aerylamide-2-methylpropanesulfonic acid was charged and neutralted (60 meN) by adding dropwise 78.4 g of an aqueous sodium hydroxide solution containing 1.9 g of addium hydroxide dissolved therein while cooling from outside. To this solution, 14.7 g of arrylamide and 23.1 mg of NN-methyleneblascrylamide were added and then 0.05 g of ammonitum persulfate was added and dissolved. The same operation was conducted, except that the resulting aqueous solution containing a polymerization inliation and a crossiningly

agent, which is essentially composed of 2-acrylamide-2-methylpropanesulfonic acid, a sodium salt thereof and acrylamide as a main component was used, a water absorbent material of the present invention as gel-like particles was obtained. The evaluation results of characteristics of the water absorbent material are shown in Table 6.

5 [Example 8]

[0144] In a 500 ml Erlenmeyer (lask, 30 g of polymer particles oblained in Example 7 were weighed, and then a mixed solution of 1.2 g of actions, 2.1 g of one exchange water, 0.0 g of tylcytly metaprate and one of the property of the prop

[Comparative Example 4]

[0145] The same operation as in Example 7 was conducted to prepare an aqueous solution containing a polymetrcation initiator and a crosslinking agent, which is essentially composed of 2-excylamide2-methylpropaeausiforinic acid, a sodium salt thereof and acrylamide as a main component, was prepared. The same operation as in Comparative Example 3 was conducted to obtain a water absorbent material of the present invention as gel-like particles. The evaluation results of the resulting water absorbent material are shown in Table 6.

[Comparative Example 5]

[0146] 4.44 g of polyethylene glycol diacrylate was dissolved in 5500 g of an aqueous solution of sodium acrylate (neutralization ratio: 75 mo/%), and after deaerating with a nitrogen gas, 2.4 g of sodium pesulitate and 0.12 g of 1-ascorbic acid wore added and the polymeration was conducted. After the completion of the polymeration, the resulting hydrous gel-like particles were further ground and dried in a hot-air dryer at 150°C so that the water content of the hydrous gel-like particles became 5% of less. The dried substance was granulated by a roll granulater and then passed through an ASTM 20 mesh metal wire to obtain a water absorbent polymer in an amorphous ground form.

[0147] 100 parts of this water absorbent polymer was mixed with an aqueous solution of 1 part of sodium polyaspartate (molecular weight: 10000) and 5 parts of water to obtain an absorber composition. The evaluation results of characteristics of this water absorbent material are shown in Table 6.

[0148] Components to be charged of Examples 6 to 8 and Comparative Examples 3 and 4 are shown in Table 5.

Table 5

		_							_	_									_
Comp.	. ardum				,	20			1		18.4	8.3	76.5	0.00039	0.05				
Comp.				-	-			. -	18.4	16.5	18.4		80.9	0.00039	0.05	1	-	-	
Example 8	3	0.3	1.2	3.2	0.75	20	0.75	164		16.5	18.4	61	76.5	0.00039	0.05	60'0	60.0	2.1	-
Example 7	3	0.3	1.2	3.2	0.75	20	0.75	164		16.5	18.4	1.9	76.5	0.00039	0.05				1
Example 6	3	0.3	1.2	3.2	0.75	20	0.75	164	18.4		18.4		80.9	0.00039	0.05				1
omponents to be charged (g)	Polysuccinimide (2)	GMA	NaOH	Ion exchange water	Sucrose ester F-160 (HLB=16)	Cyclohexane	Sucrose ester F-90 (HLB=9)	Cyclohexane	Na sulfomethyl methacrylate	AMPS	Acrylamide	NaOH	Ion exchange water	MBAA	APS	GMA	APS	Ion exchange water	A CONTRACTOR OF THE PARTY OF TH
ඊ			(3) First step	,							(3) Second step	· · · · · · · ·				(3) Third step	-		
	Example 7 Example 8 Example 3	Example 6 Example 7 Example 8 Comp. (Comp. 2) 3 3 Example 3 Example 3 Example 3 Example 3 Example 3 Example 4 Example 5 Example 6 Example 6 Example 6 Example 6 Example 7 Example 7 Example 7 Example 8 Example 8 Example 8 Example 9 Exampl	Example 6 Example 7 Example 8 Example 3 3 3 63	Components to be charged (g) Example 6 Example 7 Example 8 Components of Example 3 Polysuccinimide (2) 3 3 3 Example 3 GMA 0.3 0.3 0.3 0.3 NaOH 1.2 1.2 1.2 1.2	e(2) 3 3 6 Example 8 Example 3 6 Comp. a feature 1 2 12 12 12 12 12 12 12 12 12 12 12 12	Components to be charged (g) Example 6 Example 7 Example 8 Components of Example 3 Polysuccinimide (2) 3 3 3 Example 3 GMA 0.3 0.3 0.3 0.3 NaOH 1.2 1.2 1.2 1.2 Ion exchange water 3.2 3.2 3.2 3.2 Surrose ester F-160 (HLB-16) 0.75 0.75 0.75 0.75	Components to be charged (g) Example 6 Example 7 Example 8 Components of Example 3 Polysuccinimide (2) 3 3 3 Example 3 GMA 0.3 0.3 0.3 0.3 In a conscious consistence water 3.2 3.2 3.2 Surrose ester F-160 (HLB-16) 0.75 0.75 0.75 Cyclobexane 20 20 20	Components to be charged (g) Example 6 Example 7 Example 8 Example 8 Example 3	Components to be charged (g) Example 6 Example 7 Example 8 Components Polysuccinimide (2) 3 3 3 Example 3 BMA 0.3 0.3 0.3 0.3 0.3 In Modify 1.12 1.12 1.2 1.2 1.2 In conscious exister F-160 (HLB-16) 0.75 0.75 0.75 0.75 0.75 Sucrose ester F-50 (HLB-9) 0.75 0.75 0.75 0.75 0.75 Sucrose ester F-90 (HLB-9) 0.75 0.75 0.75 0.75 0.75 Cyclohexame 164 164 164 1.64 0.75	Components to be charged (g) Example 6 Example 7 Example 8 Comp. Polysuccinimide (2) 3 3 3 Example 3 Polysuccinimide (2) 3 3 3 Example 3 GMA 0.3 0.3 0.3 0.3 In Conscious with a constraint of the constrai	Components to be changed (g) Example 6 Example 7 Example 8 Components 3 3 Components 3 Polysaccinimide (2) 3 3 3 Example 3 GMA 0.3 0.3 0.3 0.3 In on exclude water 3.2 3.2 3.2 Surrose ester F-00 (HLB-16) 0.75 0.75 0.75 Sucrose ester F-00 (HLB-9) 0.75 0.75 0.75 Cyclohexane 164 164 164 Na sulfomethyl methacrylate 184 165 16.5 16.5 AMPS 165 16.5 16.5 16.5	Components to be charged (g) Example 6 Example 7 Example 8 Comp. Polysuccinimide (2) 3 3 3 Example 3 Polysuccinimide (2) 3 3 3 Example 3 GMA 0.3 0.3 0.3 0.3 Example 3 NuCOH 1.2	Components to be changed (g) Example 6 Example 7 Example 8 Comp. Polysuccinimide (2) 3 3 3 Example 3 GMA 0.3 0.3 0.3 6.3 NaOH 1.2 1.2 1.2 1.2 Ion exclange water 3.2 3.2 3.2 - Sucrose ester F-160 (HLB-16) 0.75 0.75 0.75 - Sucrose ester F-50 (HLB-9) 0.75 0.75 - - Sucrose ester F-50 (HLB-9) 0.75 0.75 - - Sucrose ester F-50 (HLB-9) 0.75 0.75 - - Sucrose ester F-50 (HLB-9) 1.75 1.64 1.64 - - Aviolence of the Child in the	Components to be changed (g) Example 6 Example 7 Example 8 Components to be changed (g) 3 3 3 5	Components to be changed (g) Example 6 Example 7 Example 8 Example 3	Components to be changed (g) Example 6 Example 7 Example 8 Comp. Polysuccinimide (2) 3 3 3 Example 3 GMA 0.3 0.3 0.3 6.3 MaOH 1.2 1.2 1.2 1.2 Ion exclamage water 3.2 3.2 3.2 - Sucrose ester F-160 (HLB=16) 0.75 0.75 0.75 - Sucrose ester F-90 (HLB=9) 0.75 0.75 0.75 - Sucrose ester F-90 (HLB=9) 0.75 0.75 0.75 - Cyclohexane 164 164 164 164 Na sulfometryl methacytate 184 184 184 184 Acrylamide 184 184 184 184 Ion exchange water 80.9 0.00039 0.00039 0.00039 APPS 0.05 0.05 0.05 0.05 0.05 0.05	Description of the components to be changed (g) Example 6 Example 7 Example 8 Example 3	Description of the components to be changed (g) Example 6 Example 7 Example 8 Example 3	Components to be changed (g) Example 6 Example 7 Example 8 Comp. Polysuccinimide (2) 3 3 3 Comp. Mod M 11.2 1.2 1.2 Example 3 Nature ester F-160 (HLB-16) 0.75 0.75 0.75 0.75 Sucrose ester F-160 (HLB-16) 0.75 0.75 0.75 - Sucrose ester F-160 (HLB-9) 0.75 0.75 0.75 - AMPS 164 164 164 - - ANI sulfomethyl methacylate 18.4 18.4 18.4 - - - - - - - - - - - - - - -

-	- 0.3
Acetone	Hydrophilic silica 200CF

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GMA: glycidyl methacrylate

AMPS: 2-acrylamide-2-methylpropanesulfonic acid

MBAA: N,N'-methylenebisacrylamide

APS: ammonium persulfate

Polysuccinimide (2): polysuccinimide (Mwr. 3,000) obtained in Synthetic Example 2

Table 6

	W	Water absorption ratio (g/g)	g)	Water absorption	
	Ion exchange water	0.9% aqueous solution of NaCl	Artificial seawater	rate (seconds)	Gel strength (g/cm²)
Example 6	71.5	121	55	120	2.4
Example 7	520	49	30	64	6.0
Example 8	490	45	28	55	12.0
Comp. Example 3	587	86	47	195	2.4
Comp. Example 4	440	41	2.7	95	6.0
Comp. Example 5	380	37	3	257	1.2

[0149] As is apparent from Table 6, the water absorbent materials of the present invention have high water absorption ratio and high water absorption rate in ion exchange water, an aqueous 0.9% NaCl solution, and artificial seawater as compared with the water absorbent materials of Comparative Examples and water absorption performances are improved.

INDUSTRIAL APPLICABILITY

[0150] The water absorbent material of the present invention has high water absorption properties for high concentration salt-containing solutions such as seawater, aqueous deliquescent calcium chloride solution, blood, and body fluids (e.g. urine, sweat, etc.), as compared with a conventional water absorbent material by using a copolymer containing polysaccharides as a copolymer component, and can improve water absorption properties to salt-containing solutions without impairing a water absorption ratio and a water absorption rate by using a monomer having a sulfionic acid group containing othylenically unsaturated double bond. Therefore, the water absorbtent material of the present invention can be widely used as water absorbent materials for water sealing materials, civil engineering materials, materials for agricultural/inclutural disposable sanitary materials, and household articles.

[0151] The absorbent article of the present invention is suited for use in disposable dispers for infants, adults, and persons suffering from incontinence, and in sanitary napkins because of its excellent absorbency of urine and body fullids as well as excellent urine leakage inhibition effects.

Claims

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- A water absorbent material comprising a copolymer of an anhydropolyamino acid having at least one ethylenically unsaturated double bond in a molecule (A) and a water-soluble monomer having an ethylenically unsaturated double bond (B).
 - A water absorbent material according to claim 1, wherein the water-soluble monomer having an ethylenically unsaturated double bond (8) is at least one selected from the group consisting of (meth)acrylic acid, alkall metal salt of (meth)acrylic acid, ammonium salt of (meth)acrylic acid, and an amidiated compound of (meth)acrylic acid.
 - 3. A water absorbent material according to claim 1, wherein the water-soluble monomer having an ethylenically unsaturated double bond (8) is a monomer having an ethylenically unsaturated double bond, and a sulfonic acid group and/or a sulfonate group in a molecule.
- 35 4. A water absorbent material according to claim 1, wherein the anhydropolyamino acid having at least one ethylenically unsaturated double bond in a molecule (A) is a reaction product of an anhydropolyamino acid having no ethylenically unsaturated double bond in a molecule (A-1) and a compound which has an ethylenically unsaturated double bond and a functional group having reactivity with the anhydropolyamino acid (A-1) in a molecule (A-1) and a compound which has a compound which has an ethylenically unsaturated double bond and a functional group having reactivity with the anhydropolyamino acid (A-1) in a molecule (A-1).
- A water absorbent material according to claim 1, wherein the copolymer comprises gel particles.
 - A water absorbent material according to claim 4, wherein the compound which has an ethylenically unsaturated double bond and a functional group having reactivity with the anhydropolyamino acid (A-1) in a molecule (A-2) is a compound represented by the following general formula (§):

wherein R1 represents at least one functional group selected from the group consisting of amino group, epoxy group, carboxly group, carbodimide group, oxazoline group, imino group and isocyanatia group. O represents an alkylene group having 1 to 10 carbon atoms, and R2 represents hydrogen or an alkylene group having 1 to 4 carbon atoms.

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- A water absorbent material according to claim 4, wherein the anhydropolyamino acid having no ethylenically unsaturated double bond in a molecule (A-1) is polysucclnimide.
- A water absorbent material according to claim 1, wherein a portion or all of the anhydropolyamino acid having an ethylenically unsaturated double bond in a molecule (A) is hydrolyzed.
 - A water absorbent material according to claim 1, wherein a water absorption ratio of a physiological saline solution is 10 g/g or more.
- 10. A water absorbent material comprising a copolymer of an anhydropolyamino acid having at least one ethylenically unsaturated double bond in a molecule (A), a water-soluble monomer having an ethylenically unsaturated double bond (B), and polyaccheridos (C).
 - 11. A water absorbent material according to claim 10, wherein the anhydropolyamino acid having at least one ethylenically unsaturated double bond in a molecule (A) is a reaction product of an anhydropolyamino acid having no ethylenically unsaturated double bond in a molecule (A-1) and a compound which has an ethylenically unsaturated double bond and a functional group having reactivity with the anhydropolyamino acid (A-1) in a molecule (A-2).
 - 12. A water absorbent material according to claim 10, wherein the copolymer comprises gel particles
 - 13. A water absorbent material according to claim 11, wherein the compound which has an ethylenically unsaturated double bond and a functional group having reactifyly with the anhydropolyamino acid (A-1) in a molecule (B) is a compound represented by the following general formula [i]:

wherein R¹ represents at least one functional group selected from the group consisting of amino group, epoxy group, carboxyl group, carboxilloring group, oxazoline group, imino group and isocyanate group, O represents an alkylene group having 1 to 10 carbon atoms, and R² represents hydrogen or an alkylene group having 1 to 4 carbon atoms.

- 14. A water absorbent material according to claim 11, wherein the anhydropolyamino acid having no ethylenically unsaturated double bond in a molecule (A-1) is polysuccinimide.
- 49 15. A water absorbent material according to claim 10, wherein a portion or all of the anhydropolyamino acid having an ethylenically unsaturated double bond in a molecule (A) is hydrolyzed.
 - 16. A water absorbent material according to claim 10, wherein a water absorption ratio of a physiological saline solution is 10 g/g or more.
 - 17. An absorbent article comprising an absorber comprising a water absorbent material and a fiber material arranged between a liquid-permeable sheet and a liquid-impermeable sheet, wherein the water absorbent material comprising a copolymer of an anhydrophysmino add having at least one ethylenically unsaturated double bond in a molecule (A) and a water-soluble monomer having an ethylenically unsaturated double bond (B).
 - 18. An absorbent article comprising an absorber comprising a water absorbent material and a fiber material arranged between a liquid-permeable sheet, wherein the water absorbent material so a water absorbent material comprising a copolymer of an antilydroplyamino add having at least one ethylencially unsaturated double bond in a molecule (A), a water-soluble monomer having an ethylenically unsaturated double bond (B), and polyacacharides (L).
 - 19. An absorbent article according to claim 17, wherein the copolymer comprises gel particles.

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20. An absorbent article according to claim 18, wherein the copolymer comprises gel particles.

INTERNATIONAL SEARCH REPORT

PCT /.TD01 /06428

A. CLASSIFICATION OF SUBJECT MATTER Int.Cl ⁷ CO8F290/14, AS1F13/53,									
According to International Patent Classification (IPC) or to both national classification and IPC									
B. FELD									
Minimum documentation searched (classification system followed by classification symbols) Int. Cl ² CO8F290/00-290/14, A61F13/15-13/84,									
	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched [Recursive data have coverabled during the international search (name of data base and, where executed by, such term used)								
7									
C. DOCUMENTS CONSIDERED TO BE RELEVANT									
Category*	Citation of document, with indication, where sp		Relevant to claim No.						
P,A	JP 2000-290370 A (Dainippon In) 17 October, 2000 (17.10.00), Claims (Family: none)	t and Chemicals, Inc.),	1-20						
P,A	JP 2000-281915 A (Dainippon Ink and Chemicals, Inc.), 1-20 10 October, 2000 (10.10.00), Claims (Femily: mone)								
А	JP 2000-63511 A (Mitsui Chemicals, Ltd.), 1-29 Pebruary, 2000 (29.02.00), Claims & JP 11-60729 A								
A	JP 11-158266 A (Mitsui Chemica: 15 June, 1999 (15.06.99), Claims (Family: none)	ls, Ltd.),	1~20						
A	JP 10-298282 A (Nippon Shokuba: 10 November, 1998 (10.11.98), Claims (Family: none)	Co., Ltd.),	1-20						
Purthe	r documents are listed in the continuation of Box C.	See patent family annex.							
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"L" deceme	ent which may throw doubts on priority claim(s) or which is establish the publication date of another citation or other	step when the document is taken alone "Y" document of particular relevance; the	Isimed invention cannot be						
"O" deceme	special reason (as specified) Considered to involve an inventive step when the document is combined with one or more other such documents, such								
"P" docume	monns combination being obvious to a person skilled in the art								
Date of the B	ormul completion of the international search october, 2001 (23.10.01)	Date of mailing of the international sear 06 November, 2001 (0							
	ailing address of the ISA/ nese Patent Office	Authorized officer							
Facsimile No	Facsimile No. Telephone No.								

Form PCT/ISA/210 (second sheet) (July 1992)